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METHOD OF IMPROVING GENE TRANSFER EFFICIENCY INTO PLANT CELLS

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A method for gene introduction by which higher efficiency for gene introduction than that by the conventional Agrobacterium method may be attained simply and without injuring the tissue is disclosed. According

to the method of the present invention, the efficiency of

gene introduction into plant cells by a bacterium belong ing to genus Agrobacterium is promoted by accompanying centrifugation of the plant cells or plant tissue.

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Description

Technical Field

[0001] The present invention relates to a method for promoting efficiency of gene introduction into plant cells.

Background Art

the high efficioncy, the small number of copies of the introduced gene, the feature that the gene may be introduced without fragmenting a specific region called T-DNA, and the feature that the frequency of mutation occurred during cultivation is low because transformants may be obtained by cultivation for a short period of time. Therefore, the method [0002] The method for transformation using Agrobacterium has a number of excellent features including, in genera is widely used as the most useful method for transforming various plants. 5

Although the Agrobacterium method is an extremely excellent method for transforming plants, whether the genotype and the plant itssue used (Potrykus et al. 1998 (Reference (33))). That is, there are species with which the transformation has not been successful, and species with which the transformation may be attained only with Irnited varieties. Further, there are species with which the tissue to be used is limited so that a large amount of materials cannot be treated. To prepare a practical variety by genetic recombination, it is necessary to prepare a large number of transformed plants and to select the line having the desired character therefrom. However, at present, the type of plants with which a large number of transformed plants may be prepared for this purpose is limited. Thus, to develop transformation is successful or not and the transformation efficiency largely varies depending on the plant species an improved method by which this problem may be overcome is strongly demanded. [0003] 5 8

[0004] Although the method for transformation via Agrobactertum differs in the starting material, composition of the culture medium and the like, it is almost common to the Agrobacterium methed that the method comprises making a McCormick 1991 (Reference (29)), Lindsey et al. 1991 (Reference (28))). Thus, studies for improving transformation system has been carried out mainly on the Agrobacterium strain, constitution of the vector, composition of medium tissue which is a starting material contact a suspension of *Agrobacterium*, selecting transformed cells after co-culturing and growing transformed plants. The Agrobacterium is infected without performing a special treatment except for ster ilization treatment which is carried out as required (Rogers et al. 1988 (Reference (34)), Visser 1991 (Reference (38)) 23

state in which the genes are likely to be introduced have been scarcely made. If the physiological state of the itssue [0005] On the other hand, studies for changing the plant tissue before infection of Agrobacterium to a physiological in addition to the promotion of the transformation efficiency, transformation may be attained for the species or genotypes can be changed to such a physiological state by a simple treatment, the mathod is very useful, and it is expected that types of selection marker gene and promoter, the type of the itssue used as the material, and the like. 8

with which transformation has been hitherto difficult, that is a prominent effect. Known studies about pretreatment of plant tissue include particle gun treatment (Bidney et al., 1992 (Reference (5))) and ultrasonication treatment (Trick et ically injuring the tissue, so as to increase the number of plant cells infected. However, these methods are nothing more than developments of the leaf disk method (Horsch et al., 1985 (Reference (17))) and nol treatments based on novel concepts. The degree of effectiveness and universality of the methods have not been claritied, and they are not al., 1997 (Reference (37)). Both of these methods alm at promoting invasion of bacteria into the plant itssue by physused as general methods 8 \$

Disclosure of the Invention

[0006] Accordingly, an object of the present invention is to provide a method for promoting efficiency of gene introduction into plant cells, by which gene introduction can be attained simply with a higher efficiency than the conventional gene introduction by Agrobacterium method, without injuring the tissue. ŧ,

[0007] The present inventors intensively studied to discover that in the gene introduction method using Agrobacto*rium,* the gene introduction efficiency may be significantly promoted by centrifuging the plant cells or plant tissue sub-

[0008] That is, the present invention provides a method for promoting efficiency of gene introduction into plant cells by a bacterium belonging to genus Agrobacterium, comprising contrifuging said plant colls or plant tissue. ected to the gene introduction, thereby completing the present invention. 8

[0009] By the present invention, a method for promoting efficiency of gene introduction into plant cells, by which terium method, without injuring the tissue, has been provided. The method of the present invention may be applied to gene introduction can be attained simply with a higher efficiency than the conventional gene introduction by Agrobac both monocotyledons and dicotyledons.

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Brief Description of the Drawings

Fig. 3 is a schematic view for showing the intermediate vector system and binary vector system which are major two voctor systems of bactoria bolonging to gonus Agrobactorium. Fig. 2 is a gene map of pSB133 which is an example of super-binary vectors, that may preferably be employed in Fig. 4 is a schematic view showing Iwo binary vector systems derived from super virulent strain A281 of Agrobac-Fig. 1 is a drawing for showing a method for constructing pTOK233 which is an example of super-binary vectors, that may preferably be employed in the present invention.

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[0011] In the above drawings, the following reference symbols denote the following meanings.

vids: the vid gene in the virulence region of Ti plasmid pTI80542 contained in Agrobacterium tumefactens A281 BL: left border sequence of T-DNA of bacteria belonging to genus Agrobacterium BR: right border sequence of T-DNA of bacteria belonging to genus Agrobacterium TC: tetracycline resistant gene virC: the virC gene in the virulence region of Ti plasmid pTiBo542 contained in Agrobacterium turnefacions A281 viA: the viA gene in the virulence region of Ti plasmid pTIBo542 contained in Agrobacterium tumefaciens A281 Vir. entire *vir* region of π plasmid of bacteria belonging to genus *Agrobacterium* S Vir. entire *vir* region of π plasmid ρπΒο**ς**42 of super virulent bacteria belonging to genus A*grobacterium* Tnos: terminator of nopaline synthetase gene Pnos: promoter of nopaline synthetase gene NPT, NPTII kanamycin resistant gene ORI, ori: replication origin of ColE1 SP: spectinomycin resistant gene H: restriction enzyme Hind III site HPT: hygromycin resistant gene COS, cos: COS site of \textit{\chi} phage K: restriction enzyme Kpn I site Ampr: ampicillin resistant gene P35S: CaMV 35S promoter IG: Intron GUS gene BAR: bar gene

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Best Mode for Carrying out the Invention

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s vir: tragment containing a part of vir region of TI plasmid pTIBo542

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especially 1000G to 150,000G for about 1 second to 2 hours, and the appropriate centrifugation conditions for the promoted even if the centrifugation time is very short, for example, 1 second or less. On the other hand, when the centritugal acceleration is smail, the efficiency of introducing genes may be significantly promoted by conducting the The conditions for centrilugation may appropriately be selected depending on the type of the plant used and the like, and may usually be carried out under a centrifugation acceleration of 100G to 250,000G, preferably 500G to 200,000G, more praferably 1000G to 150,000G. The time for centrifugation may appropriately be selected depending on the centrifugal acceleration, type of the plant used and so on, and is usually and preferably not less than one second. There is no upper limit of the centrifugation time, and about 10 minutes may usually be sufficient for attaining the object of the contrifugation. When the centrifugal acceleration is large, the efficiency of introducing genes may be significantly centritugation for a long time. In most cases, especially preferred centritugation conditions are about 500G to 200,000G. plant cells or tissue, or the plant cells or tissue may be contacted with the bacterium belonging to genus. Agrobacterium white contrituging the plant cells or tissue. Proforably, the plant cells or plant tissue are(is) contacted with the bacterium belonging to genus Agrobacterium, comprises centrifuging the plant cells or plant issue. The plant cells or plant tissue may be contacted with the bacterium belonging to genus Agrobacterium under normal gravity after centrifuging the [0012] The method of the present invention for promoting efficiency of gene introduction into plent cells by a bacterium belonging to genus Agrobacterium under normal gravity after centrifuging the plant cells or tissue. 23 8 ş

[0014] The method of the present invention is characterized by using the plant cells or plant tissue which were(was) particular plant cells or tissue may be easily selected by a routine experiment.

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centrifuged, or by contacting the plant cells or plant tissue with a bacterium belonging to the genus Agrobacterium while conducting centrifugation, and as the method for gene introduction or transformation *per se* using the bacterium The method for gene introduction or transformation per se into plants using a bacterium belonging to the belonging to the genus Agrobacterium, a well-known method may be appiled as it is.

the virulence region (vir region) in the TI plasmid is required for the excision of T-DNA and its transfer to plants, and the border sequences existing at the both ends of the T-DNA are necessary for the T-DNA to be exclsed. Agrobacterium thizogenes which is another bacterium belonging to the genus *Agrobacterium* has a similar system on the Ri plasmid gall disease in a number of dicotyledons. In 1970s, it was discovered that Ti plasmid concerns the vinulence, and that the T-DNA the T-DNA which is a part of Ti plasmid is incorporated into the plant genome. Thereafter, it was proved that the T-DNA the T-DNA which is a part of Ti plasmid is incorporated into the plant genome. the genes are expressed in plants in spite of the fact that the genes are bacterial genes. A group of genes existing in genus *Agrobacterium* is well-known in the art and is widely used. [0016] It is known for a long time that a soil bacterium *Agrobacterium (Agrobacterium tumefactens*) causes crown contains genes participating in synthesis of hormones (cytokinins and auxins) required for induction of tumor, and that

[0017] Since T-DNA is incorporated into the plant genome by infection of Agrobacterium, it was expected that a desired gene may be incorporated into the plant genome by inserting the desired gene in the T-DNA. However, since Tiplasmid is as large as not less than 190 to, it was difficuit to insert a gene into the T-DNA by a standard technique 5

terium may be carried out by electroporation method, triparental mating or the like). Binary vector includes pBiN19 (Bevan, 1984 (Reference (19)), pGA482 (An et al., 1988 (Reference (2)), gevan, 1984 (Reference (4)), pBi121 (Jafferson, 1987 (Reference (19))), pGA482 (An et al., 1988 (Reference (2)), Japanese Laid-open Patent Application (Kokai) No. 60-70080 (EP 120516)), and a number of new binary vectors have been constructed based on those vectors. In the system of Ri plasmid, similar vectors have been constructed and are Introduced into Agrobacterium having a disarmed type TI plasmid. The introduction of the binary vector into Agrobac-Thus, the binary vector is a small plasmid which is replicable in both Agrobacterium and E. coll, and this plasmid is ((Hoekems et al., 1983). The vir region contains virA, virB, virC, virD, virE and virG (Plant Biotechnology Encyclopodia (Enterprise Co., Ltd. (1989)), and the vir region is defined as those containing all of virA, virB, virC, virD, virE and virG. plasmids from which hormone synthetase genes were eliminated, were prepared (Fig. 3). Two methods employing such a strain, that is, a method by which a desired gene is introduced into the TI plasmid of Agrobacterfum, and a such a strain, that is, a method by which a desired gene is introduced into the TI plasmid of Agrobacterfum, and a (EP116718)). In this method, an intermediate vector which is easy to handle by genetic manipulation techniques, in which a desired gene may be inserted, and which can be replicated in E. coif is introduced into the T-DNA in the disarmed type T plasmid of Agrobacterium by triparental mating (Ditta et al., 1980 (Reference (8))). Another method is the so called binary vector method (Fig. 3), which is based on the fact that atthough the virregion is necessary for the T-DNA to be incorporated into plants, it is not necessary that the T-DNA and the wiregion exist in the same plasmid method by which a T-DNA having a desired gene is introduced into *Agrobacterium*, were developed. One of these mothods is the se called intermediate vector method (Fraley et al., 1985 (Reference (9)); Fraley et al., 1983 (Reference (10)); Zambryski et et., 1983 (Reference (40)), Japanese Laid-open Patent Application (Kokal) No. 59-140885 bryski et al., 1983 (Reference (40))), and GV3T11SE (Fraley et al., 1985 (Reference (9))), that have tumorigenic TI [0018] First, clisarmed strains such as LBAA404 (Hoekema et al., 1983 (Reference (12))), CSBC1(pGV3850) (Zamof genetic engineering. Thus, a method for introducing a foreign gene into the T-DNA was developed. 8 8 2 8

is wide and whose efficiency of transformation is higher than other strains (Hood et al., 1987(Reference (13)); Komari, 1989 (Reference (21))). This feature is brought about by a TI pissmid pTBoc42 contained in A281 (Hood et al., 1984) [0019] Agrobacterium A281 (Watson et al., 1975 (Reference (39))) is a super-virulent strain, whose host spoctrum

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at least vir8 or virG, more preferably a fragment at least containing vir8 and virG) substantially lacking at least one of the fragments of vir region is incorporated into the plasmid having the T-DNA, i.e., the binary vector. To introduce a T-DNA region into which a desired gene has been inserted into an Agrobacterium having the super-binary voctor, 1996 (Reference (25))). It has been proved that the super-binary vector gives much higher transformation efficiency than the above-described various vector systems for a number of plant species (Hiel et al., 1994 (Reference (11)): (virA, virB, virC, virD, virE and virG) (each of these may also be hereinather referred to as *vir fragment region") and a plesmid having T-DNA, this is a kind of the binary vector system. However, it is different from the binary vector in that a super-binary voctor (Komari, 1990a (Reference (22))) in which a virregion fragment (preferably a fragment containing homologous recombination via the triparental mating method may be employed as an easy method (Komari et al., (Hiei et al., 1994 (Reference (11)); Ishida et al., 1996 (Reference (18)); Komari et al., 1999 (Reference (26)). WO94/00977, WO95/06722) (Fig. 4). Since this system comprises a disarmed type TI plasmid having the vir region 1986) and EHA105 (Hood et al., 1993) containing a Ti plasmid which is a disarmed type of pTIB0542. By applying Two new systems using pTIBo542 has been developed. One system utilizes strains EHA101 (Hood at Al., these strains to the above-mentloned binary vector system, a system having a high efficiency of transformation was achieved, which is widely used for transformation of various plants. Another system is "super-binary" vector system (Reference (16)); Jin et al., 1987 (Reference (20)); Komari et al., 1988 (Reference (24))).

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shida et al., 1996 (Reference (18)); Komari, 1990b (Reference (23)); Li et al., 1996 (Reference (27)); Saito et al., 1992

stricted, and *Agrobacterium tumefactens* (e.g., the above-described A*grobacterium tumefacien*s LBA4404 (Hoekema In the method of the present invention, the host bacterium belonging to the genus Agrobacterium is not reet al., 1983 (Reference (12))) and EHA101 (Hood et al., 1986 (Reference (15))) may preferably be employed.

The method of the present invention may be applied to any of the gene introduction systems as long as it is be applied to the vector systems obtained by modification of these vectors (e.g., those wherein the entire or a part of the vir region of a bacterium belonging to the genus Agrobacterium is excised and additionally incorporated into the plasmid, or the entire or a part of the virregion of a bacterium belonging to the genus *Agrobacterium* is excised and is introduced into *Agrobacterium* as a part of a new plasmid). Further, needless to say, by the method of the present based on the expression of the group of genes in the *vir* region in the bacterium belonging to the genus *Agrobacterium* so as to obtain significant effect. Thus, the method of the present invention may be applied to any of the vector systems such as the above-described intermediate vectors, binary vectors, super-virulent binary vectors and super-binary vectors so as to obtain the advantageous effect of the present invention. The method of the present invention may also invention, the efficiency of introduction of the T-DNA region of wild type Agrobacterium is promoted so as to promote [0022]

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the desired DNA may be inserted by the triparental mating method utilizing the homologous recombination in the cell [0023] The desired gene to be introduced into the plant may be inserted into a restriction site in the T-DNA region of the above-described plasmid by a conventional method, and the Agrobacterium into which the desired gene was Incorporated may be selected based on an appropriate selection marker such as a drug resistant gene against a drug such as kanamycin or paromomycin. In cases where the plasmid is large and has a number of restriction sites, it is not always easy to insert the desired DNA into the T-DNA region by an ordinary subctoning method. In such a case, of the bacterium belonging to the genus Agrobacterium.

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[0024] Introduction of the plasmid into a bacterium belonging to the genus Agrobacterium such as Agrobacterium fumefacters may be carried out by a known method including the above-mentioned triparental mating method, elec troporation method, electroinjection method and chemical treatments with PEG or the like. ĸ

[0026] Introduction of a gene into the plant cells via a bacterium belonging to the genus Agrobacterium may be For example, a cell suspension of the bacterium belonging to the genus *Agrobacterium* having a population density of about 10⁸ to 10¹¹ cells/mi is prepared, and the plant cells or the plant tissue are(is) immersed in the suspension for only one border sequence. Alternatively, in cases where a plurality of genes are to be arranged at different sites, the plasmid may contain three or more border sequences. Alternatively, arrangement of the desired plasmid in the Ti or Al plasmid may be performed in the cell of the bacterium belonging to the genus Agrobacterium, or the desired gene attained by simply making the plant cells or plant tissue contact the bacterium belonging to the genus Agrobacterium. [0025] The gene which is to be introduced into the plant is, in principle, arranged between the left and right border sequences of the T-DNA as in the conventional method. However, since the plasmid is annular, the plasmid may contain about 3 to 10 minutes, followed by co-culturing the resultant on a solid medium for several days, thereby attaining the may be arranged in another plasmid. Further, the desired gene may be arranged in a plurality of types of plasmids. introduction of the gene.

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root, stom, fruit or any other portion of the plant. Further, dedifferentiated tissue such as a callus or a non-dedifferentiated issue such as an embryo may be employed. The type of the plant is not restricted at all, and angiosperms are preferred. [6027] The cells or the tissue to be subjected to the gene introduction are(is) not restricted at all and may be a leaf As long as the plant is an angiosperm, either dicotyledon or monocotyledon is preferred. \$

As will be concretely shown in the following Examples, by the method of the present invention, the efficiency of gene introduction is significantly promoted when compared with the conventional Agrobacterium method.

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[0029] The present invention will now be described by way of examples thereof. It should be noted that the present invention is not restricted to the following Examples.

Example 1

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(1) Agrobacterium Strains and Plasmids

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CLONETECH, U.S., (Jefferson RA 1987 (Reference (19))), LBA4404(pIG121 Hm) (Hilei, Y. et al., 1994 (Reference (11)), LBA4404(pSB133) (Fig. 2) were used.

[0031] Construction of pSB 133 was carried out as follows: A DNA fragment having a size of 6.2 kb obtained by As the Agrobacterium and its vectors, LBA4404(pB1121) (pB1121 is commercially available

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nos promoter, an hpt gene controlled by 35S promoter, and a GUS gene controlled by the 35S promoter, which GUS gene contains introns of the catalase gene of castor bean. The T-DNA region of p5B133 contains an riptil gene controlled by 36S promoter and caMV, which GUS gene contains introns of the catalase gene of castor bean (Fig. 2). The pasendis p5B133 and pTOK233 are super-binary vectors having high abilities of transformation (Koman, T. et al., 1999 (Reference (28)). This plasmid was then digested with restriction enzymes *Eco* Al and *Bgl* II to obtain a DNA fragment with a size of 8.6 kb. This DNA fragment was blunted and a Bg/ II linker (commercially available from TaKaRa) was inserted therein to of 3.1 kb containing 35S promoter and an intron-GUS gene, which fragment was obtained by digesting pIGZ21 (Ohta mosaic virus (CaMV). Each of the T-DNA regions of ptG121Hm and pTOK233 contains an nptII gene controlled by Jigesting pGA482 (An G et al., 1985 (Reference (3))) with a restriction enzyme Sal I was ligated to a DNA fragment with a size of 5.1 kbp obtained by digesting pSB11 (Kornari et al., 1996 (Reference (25)) with Sa/I to prepare a plasmid. obtain a plasmid pSB27. The pSB27 was digested with a restriction enzyme *Hind* III, and a DNA fragment with a size S et al., 1990 (Reference (32)), was inserted therein to obtain pSB33. The pSB33 was introduced into E. coll LE392. and then introduced into *Agrobacterium* LBA4404 containing pSB1 (Komarl et al., 1998 (Reference (25))) by trparental mating method (Ditta G et al., 1980 (Reference (8)). The pSB133 was obtained by homologous recombination between pSB1 and pSB33 in the cell of *Agrobacterium.* The T-DNA region of pB1121 contains a kanamycin-resistant gone (nptil) controlled by the promoter of nopaline synthetase gene (nos) and a GUS gene controlled by 35S promoter of cauliflower 5

(2) Sample Varieties and Tissues 8

[0032] As the sample varieties, Koshihikari and Tsukinohikan, which are Japonica rice varieties, were used. Glumas of immature seeds at 8 to 14 days after flowering were removed and the seeds were sterilized with 70% ethanol for several seconds and with 1% aqueous sodium hypochlorite solution containing Tween 20 for 15 minutes. After washing the scods several times with sterilized water, immature embryes with longths of 1.5 to 2 mm were excised and used as the sample tissue. æ

(3) Centrifugation Treatment

[0033] The immature embryos of rice were placed in tubes containing sterilized water and centrifuged under an acceleration of 760G to 150,000G using a micro high-speed centrifuge, large high-speed centrifuge or an utre high-speed centrifuge. After the centrifugation, the immature embryos were infected with *Agrobactentum*. 8

(4) Infection and Co-culturing

[0034] The method for infection to the immature embryos and the method for co-culturing were in accordance with the methods by Hiel et al. (1994) (Reference (11)). That is, after the centrifugation, the sterilized water in each tube was removed and suspension of A*grobactertum* was addad, followed by stirring the mixture with a vortex mixer for S 8

temperature for about 5 minutes, the immature embryos were plated on a medium for co-culturing. As the medium for co-culturing, 2N6-AS medium (Hici et al. 1994 (Reference (11))) was used except that the inorganic salts thereof were [0035] The suspensions of bacteria were prepared by collecting colonies of Agrobacterium cultured on AB medium (Chilton, M-D et al., 1974 (Reference (6))) with a platinum loop and suspending the collected bacteria in modified AA medium (AA major inorganic salts, AA amino acids and AA vitamins (Toriyama K. et al., 1985 (Reterence (38)), MS 0.2 M glucose). After leaving the mixture of immature embryos and the suspension of Agrobacterium to stand at room changed to the composition of R2 medium (Ohira et al. 1973 (Reference (31)). It should be noted, however, that the The density of the bacterial cells to be infected was adjusted to 1 x 10º to 1 x 10º cturni. The co-cuturing was carried out for 3 to 13 days, and a portion of the immature embryo was treated with X-Gluc to check the expression of the GUS gene (Hiei et al. 1994) (Reference (11)). That is, immediately after the co-culturing, the tissue was immersed in 0.1M phosphate buffer (pH 6.8) containing 0.1% Triton X-100, and was left to stand at 37°C for 1 hour. After removing Agrabacterfum with phosphate buffer, phosphate buffer containing 1.0 mM 5-bromo-4-chloro-3-Indolyl-8-D-glucuronic acid (X-gluc) and 20% methanol was added. After incubating the resultant at 37°C for 24 hours, tissueds colored in minor salts (Murashige, T et al., 1962 (Reference (30)), 1.0 g/l casamino acid, 100 µM acetosyringone, 0.2 M sucrose, major inorganic salts (KNO₃, KH₂PO₄, CaCl₂ZH₂O, MgSO₄7H₂O) were added to the modium to half concentrations. \$ ŧ, 8

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[0036] After the co-culturing, the immature embryos and call were transferred to a primary selection medium con-

taining 250 mg/l carbonicitin and 250 mg/l cefotaxime, and further containing 200 mg/l paromomycin or 10 to 30 mg/l Inggromycin, and cultured at 30°C under luminous condition for 1 to 2 weeks. As the primary selection medium, 2NBK medium described in Hiel et al. (1994) (Reference (11)) supplemented with D-sorbitol to 30 g/l was used (K medium). Further, a medium (N medium) which was the same as the 2NB medium (inorganic salts and vitamins of NB (Chu C. C. 1978 (Relierence (7))), 1 g/l casamino acid, 2 mg/l 2,4-D) except that the concentration of (NH₄)₂SO₄ was changed to 223 mg/l and that the amino acids of AA medium (Toriyama et al., 1985 (Reference (36)) were supplemented was also used in the test.

[0037] The calli formed on the primary selection medium were transferred to a secondary selection medium containing 250 mg/l cetotaxime and 250 mg/l carbenicillin, and further containing 200 mg/l paromomych or 80 mg/l hygromych. and cultured at 30°C under turninous condition for 1 to 2 weeks. As the secondary selection medium, a medium which was the same as Ne7-medium described in Hiel et al. (1994) (Relience (11)) except that the concentration of (NH₄≥SO₄ was changed to 222 mg/l and that the armino acids of AA medium (Toriyama et al., 1985 (Relience (36)) were supplemented was used. To the primary and secondary selection media containing paromomycin, agarose was added to 8 g/l as a solidifier. The rate of emerged resistant call was investigated after the secondary selection.

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(6) Regeneration of Transformants

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[0038] The call resistant to the selection drugs, obtained from the scutella of immature embryos were plated on N6S3 medtum (Hiel et al. 1994 (Reference (11)) for regeneration containing 250 mg/l carbenicillin and 250 mg/l cefo-taxine, and further containing 100 mg/l perconomycin or 50 mg/l hygromycin.

(7) Checking GUS Expression in Regenerated Plants

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[0039] Leaves of the regenerated plants resistant to the drugs, obtained by culturing for regeneration at 25°C under furninous condition for 4 to 5 weeks were checked for the expression of GUS gene by treating them with X-Gluc in the same mannor as montioned above (Hele et al. 1994 (Reference (11)). The regenerated plants were transplanted to 500-lotd diluted aqueous Hyponex solution and cultivated at 25°C under furninous condition for about 2 weeks, followed by transplantation to posts in a green house.

30 (8) Results

(i) Discussion about Effects by Centrifugation Treatment

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(1040) Using a micro high-speed centrifuge, large high-speed centrifuge and an ultra high-speed centrifuge, the efficiency of gene introduction was promoted when the acceleration was within the range of 10KG to 100KG (Tables 1, 2, 3 and 6). As for the treatment into advantageous effect was clearly observed with the treatment for 10 minutes (Tables 4, and 5). The froquency of the transiont expression of GUS was not different between the varieties, that is, between Koshihikari and Tsukhohikari Since not only the effect for promoting the efficiency of gene introduction, but also the effect for inducing idomation of callus was observed, if was suggested that centrifugation treatment is effective for induction and growth of calli and in the culturing of plants including other species.

(10041) As shown in Table 8, induction of calli from the immature embryos of Tsukinohikari was not at all observed when the centrifugation was carried out at 250KG for 60 minutes using the utra high-speed centrifuga. However, induction of calli was observed when the centrifugation was carried out at 110 KG for 60 minutes, and expression of GUS was also observed at high rate. Similarity, as for Koshilikari, induction of calli from the immature embryos of Tsukinohikari was not at all observed when the centrifugation was carried out at 250KG for 60 minutes using the utra high-speed centrifug. From those results, the advantageous effect by centrifugation for rice immature embryo is thought to be obtained at an acceleration between 5 KG to 200 KG. Thus, in view of the simplicity of the treatment, when a micro high-speed centrifuge or a large high-speed centrifuge is used, the treatment at 20KG or 40KG is thought to be appropriate. Further, as shown in Tables 9, 10 and 11, it was proved that by the centrifugation treatment at 20KG or 60 minutes, transformation using immature embryo may be attained not only for LBA4404(pSB133) having a superbinary vector known to have a high transformation ability, but also for LBA4404 (pIGT21Hm) containing an ordinary binary vector.

63 (ii) Discussion about Centrifugation Treatment and Duration of Co-culturing

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[0042] As shown in Tables 7 and 8, the efficiency of GUS expression observed in the transient assay was higher when the duration of co-culturing was 6 or 13 days than when the duration of co-culturing was 3 days. In another

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experiment, a high GUS expression was observed when the duration of co-culturing was 9 days. Various immature principos which underwent different durations of co-culturing are now cultured on a primary selection medium (10 ppm hygronycin, 200 ppm peromomycin), and there is a tendency that the rate of emerging of drug resistant call its emailer in the group co-cultured for 9 or 13 days than in the group co-cultured for 3 or 6 days.

(iii) Examination of Efficiency of Transformation by Centrifugation Treatment

[0043] At present, the GUS-positive transformants (Tables 4 and 5) prepared as described above are acclimatized, and cultuming is continued. For some lines, seeds were collected and entity was checked. As a result, no differences in morphology and fortility between the contrifuged transformants and the non-treated transformants (Koshihikari and Tisukhonihkari) were observed.

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using calli of rice. Aldemite RR et al. 1996 (Reference (1))) reported a case of transformation using rice immature may be possible to keep high efficiency of transformation by subjecting the immature embryo to the centrifugation ment method according to the present invention, even when an ordinary binary vector is used, a high efficiency of binary vector. Further, by employing both the super-binary vector and the centrifugation treatment method, the efficiency [0044] Hiel et al. (1994 (Reference (11))) reported that transformation may be attained with a relatively high efficiency embryo. To more effectively and more stably carry out these transformation methods, the above-described centrflugation treatment method is very effective. Especially, although the quality of immature embryo is likely varied depending on the environment of culturing so that it is not easy to always obtain immature embryo suited for transformation, it treatment. Hiel et al. (1994) (Reference (11)) showed that a super-binary vector having a high transformation ability promotes the efficiency of transformation of rice. According to Aldernita RR et al. 1998 (Reference (1)), transformants were obtained only in the test using LBA4404(pTOK233) containing a super-binary vector. By the centrifugation treattransformation is attained, which is comparable to or even higher than that attained in the transformation using a supermay bo oven more premoted. Still further, it is expected that transformants may be obtained by employing the centrifugation treatment method for the varieties with which a transformant has not hitherto been obtained. 2 8 23

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ifuga	3133)	ŀ
Various Centrifugation Treatments and Results of GUS Expression after Co-culturing (Sample Strain:	LBA4404/pSB133)	
SEO.	4404	
Var	Æ	

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Variety	Population Density of	Not	O	Centrifugal Acceleration	lon
	Infected Bacterium (cfu/ml)				
			760 G	8,500 G	19,100 G
Koshihikari	1×108	3/10(+)	6/10(+)	7/10(++)	7/10(+++)
	1×109	2/10(+)	0/10(-)	4/10(++)	7/10(+++)
Tsukinohikari	1×108	4/10(+)	3/10(+)	9/10(+++)	(+++)01//
	1 × 10 ⁹	1/10(+)	6/10(++)	2/10(+)	7/10(+++)

Time of Centrifugation Treatment: 10 minutes; Duration of Co-culturing: 3 to 5 days; Number of GUS-positive immature embryos/Number of sample immature embryos

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The symbols in parentheses indicate the area of the region in scutells in which GUS was expressed. ∴ none; +: small; ++: medium; +++: largo

Table 2

Rate of Emerging of Paromomycin-resistant Calli from Koshihikani Immature Embryos (Sample Strain: LBA4444/pSB133) Centrifugal Acceleration 8,500 G 760 G Not Treated Population
Density of
Infected
Bacterium
(cfu/ml) Selection Medium

Numbor of immature embryos from which resistant calli were derived/Numbor of sample immature embryos, checked

0.0%(0/19)

0.0%(0/21) 0.0%(0/22)

0.0%(0/21) 0.0%(0/23)

1×108

1×109

medium ¥

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after completion of the secondary selection Time of Centrifugation Treatment: 10 minutes; Duration of Co-culturing: 3 to 5 days

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Table 3

23	Rate of Emerging of LBA4404/pSB133)	Rate of Emerging of Peromomycin-resistant Calil from Tsukinohlikari Immature Embryos (Sample Strain: LBA4404/pSB133)	from Tsukinohike	ıri Immature Err	danyos (Sample S	Strain:
	Selection Modium	Population Density of Infoctod Bacterium (cfw/ml)	Not Treated	ő	Centrifugal Acceleration	ation
R				760 G	8,500 G	19,100 G
	z	1 x 10 ⁸	0.0%(0/11)	0.0%(0/11) 0.0%(0/11)	30.0%(3/10)	38.4%(4/11)
	medium	1 x 109	0.0%(0/11)	0.0%(0/11) 9.1%(1/11)	27.3%(3/11)	54.5%(6/11)
£	×	1 × 108	0.0%(0/10)	0.0%(0/15)	9.1%(1/11)	9.1%(1/11)
	magem	1 x 109	0.0%(0/11)	0.0%(0/11)	0.0%(0/11)	45.5%(5/11)

Number of immature embryos from which resistant calli were derived/Number of sample immature embryos, checked after completion of the secondary selection Time of Centrifugation Treatment: 10 minutes; Duration of Co-culturing: 3 to 5 days

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Table 4

Strains and Plasmids Not Time of Centrifugation Treatment Treated	Not Treated	Time of (Time of Centrifugation Treatment	reatment
		10 minutes	30 minutes 60 minutes	60 minutes
LBA4404(pSB133)	9/10(+)	9/10(++)	10/10(++)	10/10(+++)
LBA4404(pTOK233)	9/10(+)	10/10(++)	10/10(++)	10/10(+++)

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Centrifugal acceleration: 20,000G; Sample Variety: Koshihikari;

Number of GUS-positivo immature embryos/Number of sample immature embryos area of the region in scutella in which GUS was expressed. +: small; ++: medium; +++: large

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Table 5

Time of Centrifugation Treatment and Rate of Emerging of

	Paromormycin-resist	Paromormycin-resistant Calli (Variety: Koshihikari)	ihikarl)				
,	Selection Medium	Selection Medium Culturing Condition Not Treated	Not Treated	Тте об	Time of Centrifugation Treatment	eatment	
				10 minutos	30 minutos	60 minutos	
9	N medium	Luminous (30°C)	0.0%(0/31)	34.3%(12/35)	0.0%(0/31) 34.3%(12/35) 35.0%(14/40) 53.3%(16/30)	53.3%(16/30)	
		Dark (30°C)	0.0%(0/32)	54.1%(20/37)	0.0%(0/32) 54.1%(20/37) 34.2%(13/38)	58.6%(17/29)	
5	madium	Luminous (30°C)	0.0%(0/31)	20.0%(7/35)	0.0%(0/31) 20.0%(7/35) 38.5%(15/39)	40.0%(12/30)	
!		Dark (30°C)	0.0%(0/32)	48.6%(17/35)	0.0%(0/32) 48.6%(17/35) 41.0%(16/39)	33.3%(10/30)	

31.8%(7/22) 13.3%(2/15) 18.2%(4/22) 0.0%(0/22)

15.0%(3/20)

0.0%(0/22) 4.5%(1/22)

4.8%(1/21) 4.3%(1/23)

1×108

N medium

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1×109

16.7%(3/18) 14.3%(3/21) Contrifugal acceleration: 20,000G; Duration of Co-culturing: 3 to 5 days, chocked after completion of the secondary

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g

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Number of immature embryos from which resistant calli were derived/Number of sample immature embryos

Table 6

	labre o				
tensity of Centrifugation Te	Intensity of Centrifugation Treatment and GUS Expression after Co-culturing (variety: Tsukinohikari)	lon after Co	-culturing (v	rarlety: Tsu)	kinohikari)
Centrifugation Treatments	Duration of Co-culturing	Nur	nber of Imn	Number of Immature Embryos	yos
		Frequenc	y of GUS E	Frequency of GUS Expression in Scutella	n Scutella
			#	+	‡
Not Treated	3 days	9	4	0	•
	8 days	0	2	9	8
20KG ¹⁾	3 days	۰	0	2	8
	8 days	۰	0	2	80
40KG ²⁾	3 days	-	0	-	8
	6 days	•	0	0	9
110KG ³⁾	3 days	-	0	5	4
	8 days	•	0	2	80
250KG ³⁾	3 days	10	0	0	0
	6 days	5	0	0	0

Sample Strain: LBA4404/plG1214m; Time of Centrifugation Treatment: 60 minutes 1) micro high-speed centrifuge; 2) large high-speed centrifuge 3) ultra high-speed centrifuge Rate of GUS-expressed area in scutella: : none; ±: <1/6: +: 1/8-1/4; ++: >1/4 Ş

Table 7

	Centrilugation Treatment, Duration of Co-cutturing and GUS Expression after Co-cutturing (Variety: Tsukinohikari)	ion of Co-culturing and GUS Ex	pression afte	r Co-culturing	(Variety: Ter	ukinohikari)
99	Centrifugation Treatments	Duration of Co-culturing	ž	Number of Immature Embryos	ature Embry	80
			Frequer	Frequency of GUS Expression in Scutella	xpression in	Scutella
				Ŧ	+	+
Ş		3 days	2	4	-	0
3	Not Treated	6 days	•	9	2	7
		13 davs	c	ur.	`	۳.

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Line bile on	KINDEHRAIN	S	Scutella	‡	1	n	ø	"	,	8	8	4	
	(Variety: 150	ature Embryo	opression in 9	+		S		, ,	,	7	œ		,
	r Co-culturing	Number of Immature Embryos	Frequency of GUS Expression in Scutella	,	•		,	-	-	-		•	-
ē	ression after	₹	Frequen				· •	0	0		•	5	٥
Table 7 (continued)	in of Co-culturing and GUS Exp	Duration of Co-culturing					3 days	6 days	13 days		3 days	6 days	13 days
	Continuation Treatment, Duration of Co-culturing and GUS Expression after Co-culturing (Variety: Isuking in Augustian)	Celluluganon	Centritugation Ireamierius					. Oxec	. SANS			40KG 2	

Sample Strain: LBA4404plG121Hm; 1) micro high-speed centrifuge; 2) large high-speed centrifuge; Centrifugation was carried out for 60 minutes at the indicated revolution.

Rate of GUS-expressed area in scutella: .: none; ±: <1/8; +: 1/8; +1: >1/4

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Table B

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					_	_	_	_	_	_	_	_			٦
		yos	Scutella	‡	٥			-	6	æ	6	۵	9	6	
		Number of Immature Embryos	ression in	+				7	-	2	-	4	. c		
		ser of Imme	Frequency of GUS Expression in Scutella	#	ľ	, ,	-	60	0	c			, c		,
	11	Nem	Frequency	-		_	m	-	6	, ,	, -	,	- ‹	۰ د	>
0 9091	ration of Co-culturing and C g (Variety: Koshihikari)	Duration of Co-culturing				3 days	6 days	13 days		3 days	6 days	13 days	3 days	6 days	13 days
	Centrilugation Treatment, Duration of Co-culturing and GUS Expression after Co-cutturing (Variety: Koshihikari)	Centrifugation	Treatments				Not Treated			i	20KG ¹⁾			40KG 29	

g

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Sample Strain: LBA4404plG121Hm; 1) micro high-speed centrifuge; 2) large high-speed centrifuge; Centrifugation was carried out for 60 minutes at the indicated revolution. 9

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Rate of GUS-expressed area in scutella: -: none; ±: <1/8; +: 1/8-1/4; ++: >1/4

Table 9

٠		+IGe/ACLASC .	21) Wariety Tsukinohika	Ē		
	Results of Transforms	ation by LBA4404(pp.	Results of Transformation by LBA4404(ppl 2.) (transformation by L	Mumber of GIIS.	Transformation	
	Treatments	Number of	Number of Number of Acclimatized Plants	positive Plants	Efficiency	
		ושונושוחום ביווח אמ		65	24.0(%)	
Ş	Part Transfer	50	17	71		
	NOI HEBRE			13	36.0(%)	
	Centrilugation	150	9	5		
	Treetment	_				_
	ופשחיופונו	12.00	Tage: Duration of Co-Cul	turing: 5 days		_
	Centrifugation Tr	reatment: 20KG-b0 mil	Centrifugation Treatment: 20KG-60 minutes, Duramon			

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Table 10

	Transformation	Efficiency	7.5(%)		10.6(%)		
hikarl)	Mumber of GUS.		6	,	s.	,	
Taylorikar)		Number of Number of Arctimatized Plants		6		9	
Tolanda And A	tion by LBAMMONDE	Number of	Immature Cition you	40		47	
	Results of Transforma	Treatments		11 12 12	Not Ireated	Centrifugation	Treatment
٠							ş

Centrifugation Treatment: 20KG-60 minutes; Duration of Co-culturing: 5 days

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Table 11

Transformation	Efficiency	4.1(%)	1,0,00	(w)A.A		
9170	Number of GUS- positive Plants	2		27		
Results of Transformation by LBA4404(pBi121) (Varioty: Koshihikari)	Number of Number of Acclimatized Plants		4	35	3	
tion by LBA4404(pBI12	Number of	IIIIIIIIII Cilici	64		274	
Results of Transforme	Treatments		Not Treated	101	Centrifugation	Treatment
					8	

Centrifugation Treatment: 20KG-60 minutes; Duration of Co-culturing: 5 days

Table 12

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	Transformation	Efficiency	(%)00	(2)	8.2(%)			
(Mumber of GUS.	positive Plants			30	8		
BAAAnd(nSB133) (Variety: Koshinikari)		Number of Number	Accountant of	•		ຂ		
TRAAAOA(nSB1	uoil by Edition	Number of	Immature Emoryus	64	3	281		
	Results of Transforma	Treatments			Not Treated	a discontinue	Centringation	Treatment
-								

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Centrifugation Treatment: 20KG-60 minutes; Duration of Co-culturing: 3 days

Example 2

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to centrifugation tubes containing the same medium, followed by centrifugation treatment at 20 KG or 40 KG at 4°C. the liquid medium was removed, and a suspension of LBA4404(pSB131) with a population density of about 1 imes 10 3 ctu/mi was added, followed by gently stirring of the mixture. After leaving the mixture to stand at room temperature for 5 minutes, the embryos were plated on LS-AS medium containing 10 µM AgNO₃ such that the surface of each hypocoby¹ contacts the medium. After co-culturing in the dark at 25°C for 3 days, an aliquot of the immature embryos was sampled and the transient expression of the GUS gene was checked by the treatment with X-gluc as in Example 1. The abovefor 30 minutes. After the treatment, the mixture was gently stirred and plated on LS-AS medium such that the surface of hypocotyl contacts the medium. On the other hand, infection to immature embryos after centritugation treatment of hypocotyl contacts the medium. On the other hand, infection to immature ombryos after centritugation treatment was carried out as follows: Embryos aseptically collected were washed once with LS-infilquid medium and transferred for 1 hour. The control sample was left to stand in the liquid medium at room temperature for 1 hour. After the treatment, Agrobiological Resources, The Ministry of Agriculture, Forestry and Fisheries) were aseptically collected and washed once with LS-firl liquid medium. To a centrifugal tube containing the immature embryo and 2.0 ml of LS-firl medium once with LS-firl liquid medium. To a (Reference (18))) was added to a population density of about 1 x 10° ctu/ml, and the resulting mixture was centrifuged Immature embryos of maize with a size of about 1.2 mm (variety: A188, obtained from National Institute of containing 100 µM acetosyringone, a suspension of Agrobacterium (umelaciens LBAA404(pSB131) (lahida et al. 1996 et 40,000G, at 4°C for 30 minutes. The control embryo was left to stand in the same cell suspension at room temperature S **\$ \$**

[0046] The transient expression of the GUS gene in the A188 immature embryos infected with LBA4404(pSB131) is shown in Table 13. Although any embryo showed expression of the GUS gene, a number of the immature embryos described medium and method for culturing were in accordance with Ishida, Y. et al. 1996 (Reference (18)). 8

subjected to the centrifugation treatment showed expression in larger area than the control immature embryos. The ugation treatment was performed together with the *Agrobacterium* and wherein the *Agrobacterium* was infected after ncrease in the gene-introduced sites by the centrifugation treatment was observed in both cases wherein the centrifthe centrifugation treatment. Further, expression of the GUS gene was observed in larger area than in the control even if the intensity of centrifugation and the time of treatment were changed.

possibility that the maize varieties (ishida et al. 1996 (Reference (18))) other than A188, which could not be hitherto (0047) By these results, the possibility that by culturing the immature embryos after centrifugation treatment on a selection medium, transformed plants are obtained at higher efficiency than the control was suggested. Further, the transformed by the conventional Agrobacterium method, may be transformed by the centrifugation treatment, was suggested.

Table 13

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		ransient	Translent Expression of GUS Gene in A188 Immature Embryos	re Emb	yos		
Test	Treatment	nent	Number of Sample Immature Embryos		Expression of GUS Gene	f GUS	Gene
	KG	nin		‡	‡	+	
-	40	оc	27	^	9	9	0
	Control	30	30	-	17	5	0
5	40	09	20	٥	က	17	0
	ಜ	99	82	0	9	2	0
	Control	8	8	0	-	19	0

Control was treated under 1G

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In Test 1, centrifugation treatment was performed in the presence of Agrobacterium.

In Test 2, Agrobacterium was infected after the centrifugation treatment.

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Claims

- A method for promoting efficiency of gene introduction into plant cells by a bacterium belonging to genus Agrobacterium, comprising centrituging said plant cells or plant tissue.
- The method according to claim 1, wherein said gene introduction is carried out after centrifuging said plant cells or plant tissue.
- The method according to claim 1 or 2, wherein the centritugation is carried out under a centrifugal acceleration of 100G to 250,000G.

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- The mathod according to claim 3, wherein said centrifugation is carried out under a centrifugal acceleration of 500G to 200,000G.
- The method according to claim 4, wherein said centrifugation is carried out under a centrifugal acceleration of 1000G to 150,000G.

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- 6. The method according to any one of claims 1 to 5, wherein said centrifugation is carried out for 1 second to 4 hours.
- 20 7. The method according to claim 6, wherein said centrifugation is carried out for 5 minutes to 2 hours.
- 8. A method for preparing a plant characterized by using the method according to claim 1 to 7.
- 9. Plant colls, plant tissue or plant propared by the method according to claim 1 to 8.

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- The mothod according to any one of claims 1 to 7, wherein said plant cells or plant tissue used are(is) originated from an angiosperm.
- 11. A method for preparing an angiosperm characterized by using the method according to claim 9.

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- Angiosperm cells, angiosperm tissue or angiosperm prepared by the method according to claim 10 or 11.
- 13. The method according to claim 10, wherein said plant cells or plant tissue used are(is) originated from a mono-
- A method for preparing a monocotyledon characterized by using the method according to claim 11.

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- 15. The monocotyledon colls, monocotyledon tissue or monocotyledon prepared by the method according to claim 13
- The method according to claim 13, wherein said plant cells or plant tissue are(is) originated from a plant belonging to family Gramineae.
- 17. A method for preparing a plant belonging to family Gramineae characterized by using the method according to claim 13

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- 18. The cells of the plant belonging to family Gramineae, the tissue of the plant belonging to family Gramineae, or the plant belonging to family Gramineae prepared by the method eccording to claim 18 or 17.
- 19. The method according to claim 16, wherein said plant cells or plant tissue are(is) of rice or maize.

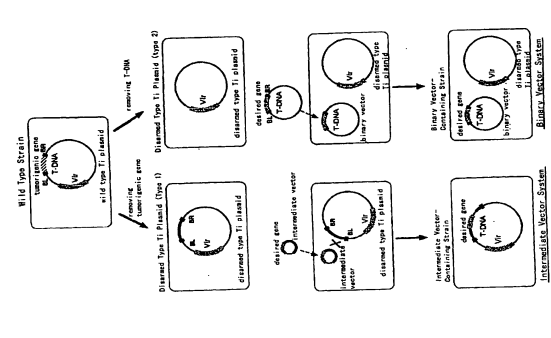
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- 20. A method for preparing rice or maize characterized by using the method according to claim 19.
- 21. Rice cells, rice tissue, rice, maize cells, maize tissue or maize prepared by the method according to claim 19 or 20.

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Кри 0.0 | Modil 0.2 | Вапн 1.2.4 | Вапн 4.3



Salt 14.9 Hndfil 15.0

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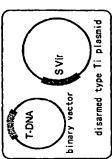
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Fig. 3

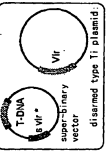
18

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Binary Vector-Containing Super-Virulent Strain



Super-Binary Vector-Containing Strain



Super-Binary Vector System

Binary Vector System by

Super-Virulent Strain

Purther documents are listed in the continuation of Box C.

F.

	INTERNATIONAL SEARCH REPORT	International application No. PCT/JP00/0	ional application No. PCT/JP00/05213
A. CLASS Int. According to	A. CLASSERCATION OF SUBJECT MATTER Int.Cl. CL2N 15/04, 5/14, MO1H 1/00, 5/00 Int.Cl. CL2N 15/04, 5/14, MO1H 1/00, 5/00 Axoodia to intrastical Parts Chaiffeation (PC) or to both material chaiffeation and PC	ں	-
B. FIELDS	FIELDS SEARCHED		
Minimun & Int.	Minimum decumentation perchod (clessification system followed by clessification symbols). Int. C1. ³ C11N 15/00-15/90, A01H 1/00-15/00		
Documental	Documentation warehol other than minimum documentation to the extent the rach documents are included in the fibrile searched	s are included i	n the fields searched
Electronic d WP I /	Pacronic dan bas occasival dring the therminant search (sears of dan leas and, when packtable, search terms seat) RP1/L (DIALCO) , BICGIS (DIALCO) , NEDLISB, JICGT PILB (JDIS)	rncikable, sean (0)	ch term uned)
C. DOCU	DOCUMENTS CONSIDERED TO BE RELEVANT		
Cettgery*	Chaden of document, with indication, where appropriate, of the relevant parages	naga.	Relevant to claim No.
۲	JP 10-179174 A (Hokko Chemical Industry Co., L 07 July, 1998 (07.07.98) (Pamily: none)	Led.),	1-21
4	WO 00/37653 A2 (THE SANJEL ROBERTS NOSLE POUNDATION, 1DKC.), 29 June, 2000 (29.06.00) 6 AU 200025943 A	ATION,	1-21
~	TRICK, H. N. et al., 'SAAT: scmication-sesisted Agrobacterius-mediated Trasformation', Transpanic Research, (September, 1997), Vol.6, No.5, pages 329-336	, No.5,	1-21
4	HORECE, R.B. et al., "A Simple and General Wethod for Transferring Genes into Plante", Science, (08 August, 1997), Vol.227, Bo.4691, pages 1229-1231	shod for 9-1231	1-1

orn: PCT/ISA/210 (second about) (July 1992)

me and mailing address of the 18A/ Japanese Patent Office

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Detr of mailing of the international search report 31 October, 2000 (31.10.00)

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